

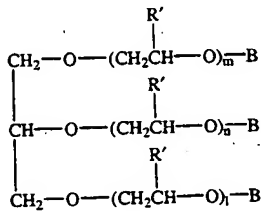
COMPOSITION COMPRISING A MIXTURE
OF ALKOXYLATED MONO-, DI- AND
TRIGLYCERIDES AND GLYCERINE

DESCRIPTION

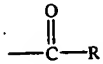
The present invention relates to a composition comprising a mixture of alkoxylated mono-, di-, and triglycerides and glycerine, to methods for the preparation of this composition, to detergent compositions comprising this composition, and to the use of the composition as surfactant or co-surfactant in detergent compositions.

Most of the known detergent compositions use anionic, amphoteric and/or non-ionic surfactants to obtain a final product showing satisfactory properties in terms of detergency and foam profile. However, most of these compositions are generally not satisfactory regarding the problem of ecotoxicity and the irritation to the eyes and the skin.

EP 0 586 323 B1 discloses detergent compositions showing improved properties regarding the ecotoxicity and the irritation to the eyes and to the skin. These compositions comprise the mono-, di- and tri-ester compounds represented by the following formula, wherein the weight ratio of mono-, di-, and tri-ester is 46-90/9-30/1-15:



wherein R' represents H or CH₃, B represents H or



wherein R represents an alkyl or alkenyl group having 6 to 22 carbon atoms, and each of m, n, and l may have a value between 0 to 40, the sum of m, n and l being in the range of from 2 to 100.

The viscosity of compositions disclosed in EP 0 586 323 B1 having a good foaming power is generally low. Although the viscosity may be increased when the alkoxylation degree is lowered, this is generally not preferred, since then the foaming power is also dramatically decreased. Therefore, a salt such as sodium chloride is generally added in order to increase the viscosity. However, adding a salt leads to an enhanced irritation of the skin and the eyes.

In view of this prior art it was the problem underlying the present invention to provide compositions showing a high viscosity and good foam stability, while also showing the good properties with respect to biodegradability and irritation to the eyes and the skin.

This problem is surprisingly solved by a composition comprising

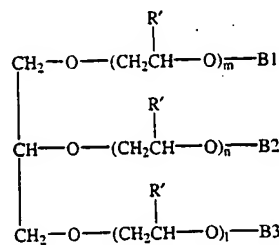
- (i) compounds represented by the following formula (I), wherein each of B1, B2 and B3 independently represent a group represented by the following formula (II);
- (ii) compounds represented by the following formula (I), wherein two of B1, B2 and B3 independently represent a group represented by the following formula (II), the remainder representing H;

(iii) compounds represented by the following formula (I), wherein one of B₁, B₂ and B₃ represents a group represented by the following formula (II); the remainder representing H;

5 (iv) compounds represented by the following formula (I), wherein each of B₁, B₂ and B₃ represent H; the weight ratio of the compounds (i)/(ii)/(iii) being 46 to 90/9 to 35/1 to 15:

Formula (I)

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R' representing H or CH₃, and each of m, n, and l independently representing a number from 0 to 4, the sum of m, n and l being in the range of 1 to 4;

Formula (II):

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30 wherein R represents an alkyl or alkenyl group having 6 to 22 carbon atoms:

The weight ratio of the compounds (i)/(ii)/(iii) in the composition of the present invention is preferably 60 to 83/16 to 35/1 to 6.

35 Particularly preferred are compounds of formula (I) wherein R' in formula (I) represents H, that is, the compounds are ethoxylated derivatives.

The sum of m, n and l in formula (I) is in the range of 1 to 4, preferably 1.5 to 3.0, more preferably in the range of 1.5 to smaller than 2.

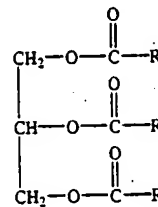
40 The weight ratio (i)+(ii)+(iii)/(iv) is preferably in the range of 85/15 to 40/60, more preferably in the range 80/20 to 45/55.

45 The compositions of the present invention can be prepared by a first method comprising the following steps:

a) Subjecting a mixture of glycerine and a compound of the following formula (III) to an interesterification reaction:

50

(III)



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60 wherein R represents an alkyl or alkenyl group having 6 to 22 carbon atoms, and

b) subjecting the reaction mixture obtained in step a) to an alkoxylation using an alkylene oxide having 2 or 3 carbon atoms in the presence of an alkaline catalyst.

65 The interesterification reaction in step a) is governed by statistics. Consequently, the molar ratio of the compounds

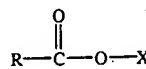
(i), (ii), (iii), and (iv) in the final product is determined by the ratio of the starting materials glycerine and the compound of formula (III). The subsequent alkoxylation reaction of step b) is a reaction which generally proceeds quantitatively, so that the amount of alkylene oxide used determines the alkoxylation degree (that is, the sum of m, n, and l). The molar ratio of the compounds (i), (ii), (iii), and (iv) is not affected by the alkoxylation, since the alkylene oxide only reacts with the remaining free hydroxyl groups in the mono- and di-ester molecules and the glycerine. However, the weight ratio of the compounds (i), (ii), (iii), and (iv) is consequently changed. Since the outcome of both reaction steps a) and b) can be predicted by the skilled person, modelling calculations can be employed to determine the correct ratio of the starting materials for a specific predetermined weight ratio of the compounds (i), (ii), (iii), and (iv) and a specific predetermined alkoxylation degree.

The compound of formula (III) includes natural fat and oil as well as synthetic triglycerides. Preferred is a fat or oil including vegetable oil such as coconut oil; palm oil; palm kernel oil; sunflower oil; rape seed oil; castor oil; olive oil; soybean oil; and animal fat such as tallow, bone oil; fish oil; hardened oils and semihardened oils thereof, and mixtures thereof. Particularly preferred are coconut oil, palm oil and tallow such as beef tallow.

Further, the composition of the present invention can be produced by a second method comprising the following steps:

- a) Reacting a mixture of glycerine and alkylene oxide having 2 or 3 carbon atoms in the presence of an alkaline catalyst.

b) Reacting the reaction mixture obtained in step a') with a compound of the following formula (IV).



wherein R is defined as above for formula (III) and X 40 represents a methyl group or H.

The degree of alkoxylation in the final product (that is, the sum of m, n, and l) is determined by the amount of alkylene oxide employed in step a'). Step b') then determines the molar ratio and the weight ratio of the compounds (i), (ii), (iii), and (iv). Again, the outcome of both reaction steps a') and b') can be predicted by the skilled person, so that modelling calculations can be employed to determine the correct ratio of the starting materials for a specific predetermined weight ratio of the compounds (i), (ii), (iii), and (iv) and a specific predetermined alkoxylation degree.

The compound of formula (IV) is preferably derived from one of the fats or oils which are preferably used in the first method of the present invention and which are listed above. Particularly preferred are tallow fatty acid and coconut oil fatty acid, palm oil fatty acid, or a methyl ester thereof.

The composition of the present invention is preferably used as a surfactant or co-surfactant in detergent compositions in which they are preferably contained in an amount of from 0,5 to 20 wt. %, more preferably 1 to 8 wt. %.

The detergent compositions of the present invention may additionally contain one or more of the following additives, depending on the purpose of the detergent composition, this list being non-limiting.

1. Anionic surfactants such as sodium alkyl ether sulphate, ammonium alkyl ether sulphate, triethanolamine alkyl ether sulphate, sodium alkyl sulphate, ammonium

alkyl sulphate, triethanolamine alkyl sulphate, sodium alpha-olefin sulphonate, sodium alkyl sulphonate, sulphosuccinates, and sulphosuccinamates.

2. Fatty acids or soaps derived from natural or synthetic sources such as coco, oleic, soya and tallow fatty acids.
3. Ethoxylated alcohols.
4. Esters of fatty acids from natural or synthetic sources such as glycol, ethylene glycol, diethylene glycol, propylene glycol, dipropylene glycol, saccharose, glucose or polyglycerine.
5. Ethoxylated fatty esters from fatty acids of hydroxyl fatty acids.
6. Amphoteric surfactants such as alkyl amidopropyl betaine, alkyl betaine, alkyl amidopropyl sulphobetaine, alkyl sulphobetaine, cocoamphoacetates, and cocoamphodiacetates.
7. Amine oxides such as dimethyl alkylamine oxides or alkyl amidopropylamine oxides.
8. Amides such as monoethanolamides, diethanolamides, ethoxylated amides or alkylisopropanolamides.
9. Alkylpolyglycosides.
10. Ether carboxylates from alcohols, ethoxylated fatty alcohols.
11. Cationic surfactants such as dialkyl dimethyl ammonium halides, alkyl benzyl dimethyl ammonium halides, alkyl trimethyl ammonium halides, esterquats derived from triethanolamine, methyldiethanolamine, dimethylaminopropanediol and oligomers of such esterquats.
12. Additives to improve such formulations, such as thickeners, pearlizing agents, opacifiers, antioxidants, preservatives, colorants or perfumes.

EXAMPLES

Compositions of the present invention were prepared according to the following methods; the values for the indicated parameters X, X', s, m, m', n, n', Y, Y', Z, Z' are shown in tables I and II:

Method 1: From Triglyceride

X g (X' moles) of triglyceride (coconut oil or palm oil), m (m' moles) of glycerine and s g of KOH 85% as catalyst are placed in a 2 kg flask properly equipped. The system is purged several times with nitrogen, vacuum stripping is carried out until 110° C., and heating is continued to 140° C. When the temperature reaches 140° C. the reactor is pressurised to 2-3 Kg/cm² with ethylene oxide added until a total of n g (n' moles).

Method 2: From Methyl Ester

m g (m' moles) of glycerine and s g KOH 85% as catalyst are placed in a 2 Kg flask properly equipped. The system is purged several times with nitrogen, vacuum stripping is carried out until 110° C. and heating is continued to 140° C. When the temperature reaches 140° C., the reactor is pressurised to 2-3 Kg/cm² with ethylene oxide added until a total of n g (n' moles). After the final charges of ethylene oxide, the reaction mixture is allowed to react for about ½ hour, z g (z' moles) of a methyl ester of fatty acid (either coconut oil fatty acid or palmoil fatty acid), is added and mixed for 45 minutes. Finally, the product is cooled and discharged from the reactor.

Method 3: From Fatty Acid

m g (m' moles) of glycerine and s g KOH 85% as catalyst are placed in a 2 Kg flask properly equipped. The system is purged several times with nitrogen, vacuum stripping is carried out until 110° C. and heating is continued to 140° C. When the temperature reaches 140° C., the reactor is pressurised to 2-3 Kg/cm² with ethylene oxide added until a total of n g (n' moles). After the final charges of ethylene

oxide, the reaction mixture is allowed to react for about $\frac{1}{2}$ hour, y g (y' moles) of a fatty acid (either coconut oil fatty acid or palm oil fatty acid), is added and mixed for 45 minutes. Finally, the product is cooled and discharged from the reactor.

The weight ratios of the mono-, di-, and triglycerides obtained by the above methods is also indicated in Tables I and II.

Then, detergent compositions were prepared with the composition of the present invention in an amount of 5 wt. % and sodium laurylethersulphate in an amount of 15 wt. %,

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the balance being water. Sodium chloride was added in the amount indicated in Tables I and II (in wt. %).

The viscosity of the compositions was then measured with a Brookfield viscosimeter at 20° C. For each experiment, a viscosity curve was prepared in order to determine the maximum (values given in cps).

The foam ability was measured at 5 seconds with a Ross-Miles apparatus using water at a temperature of 20° C. and a hardness of 20° HF. (values given in millimeters height).

The results are summarized in Tables I and II.

TABLE I

Examples according to the present inventionMixtures of 15% Active Matter of Sodium Lauryl ether Sulphate + 5% product

EXAMPLES	A	A'	B	D	E	E'	F	F'	G	H
<u>Comp.</u>										
Mono	69	69	69	77	70	70	77	77	71	78
Di	28	28	28	22	27	27	21	21	26	20
Tri	3	3	3	2	3	3	2	2	3	2
Alkyl chain (R)	Coco	Palm	Coco	Coco	Coco	Palm	Coco	Palm	Coco	Coco
EO	1,88	1,88	1,76	1,4	2,5	2,5	2,5	2,5	3,5	3,5
Prep. way										
Method	1	3	1	3	1	1	3	1	3	3
Triglyceride (x)	461,8		470,3		422,3	476,4		400		
moles trig (x')	0,69		0,7		0,63	0,57		0,48		
Fatty Acid (y)		494,3		396			329,8		351	286,2
moles FA (y')		1,85		1,87			1,56		1,66	1,35
Methyl-ester (z)										
moles ME (z')										
Glycerine (m)	252,1	283,8	256,8	382	230,6	209	318	248,5	253,9	276
moles gly (m')	2,74	3,09	2,79	4,15	2,51	2,27	3,46	2,7	2,76	3
Ethylene Oxide (n)	283,4	255,2	270,2	255,7	344,6	312,3	380,2	349,6	424,9	462
moles (n')	6,44	5,8	6,14	5,81	7,83	7,1	8,64	7,95	9,66	10,5
KOH (85%) (s)	2,7	1,2	2,7	1,2	2,4	2,2	1	1,8	1	0,8
Max visc.	45000	80000	43000	15000	22000	35000	18000	25000	26000	36000
Salt nec.	2,5	2,5	2,5	6	3,5	3,5	4	4	3,5	4
Foam	175	170	175	175	170	170	175	170	170	180

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TABLE II

Comparative ExamplesMixtures of 15% Active Matter of Sodium Lauryl ether Sulphate + 5% product

EXAMPLES	C	I	J	K	L	M	N	O
<u>Comp.</u>								
Mono	90	40	59	77	46	79	69	57
Di	10	46	35	21	42	19	28	36
Tri	0	17	6	2	12	1	3	7
Alkyl chain (R)	Coco	Tallow						
EO	2,2	2,2	4,4	0,8	3,5	4,4	15	9,9

TABLE II-continued

Comparative Examples.

Mixtures of 15% Active Matter of Sodium Laurylether Sulphate + 5% product

EXAMPLES	C	I	J	K	L	M	N	O
<u>Prep. way</u>								
Method	1	1	1	1	1	1	1	2
Tri-glycerid (x)	197,2	670,3	422,8	470	561,2	266,1	187,5	
moles trig (x')	0,29	0,99	0,63	0,7	0,83	0,39	0,28	
Fatty Acid (y)								
moles FA (y')								
Methyl-ester (z)						369		
moles ME (z')						1,3		
Glycerine (m)	376,8	111,8	146	363,6	114,9	211,3	76,8	119,3
moles gly (m')	4,1	1,22	1,59	3,95	1,25	2,3	0,83	1,3
Ethylene Oxide (n)	424,8	213,9	428,7	163,7	320,6	521,1	734,6	564,7
moles (n')	9,65	4,86	9,74	3,72	7,29	11,84	16,7	12,83
KOH (85%) (s)	1,2	3,7	2,6	2,4	3,1	1,5	1,1	1,6
Max visc.	4000	6000	13000	8000	7000	9000	5000	4000
Salt nec.	5	6	3	4	5	4	4	4
Foam	170	175	175	130	160	155	145	120

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As may be derived from the results above, when the ethoxylation degree is larger than 4 (Ex. J, M, N, O), maximum viscosity is always lower than 14000 cps measured with a Brookfield viscosimeter at 20° C. When the ethoxylation degree is lower than 1 (Ex. K) viscosity is also lower than 14000 cps. When the triester content is lower than 1 (Ex. C; 90/10/0), maximum viscosity is also very low (lower than 14000 cps). When the diester content is too high (Ex I: 40/46/17 and Ex L: 46/42/12), then the viscosity is also lower than 14000 cps.

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However, when the samples are within the alkoxylation degree in accordance with the present invention (1 to 4 EQ mols), viscosities are considerably higher (see Table I). Specially this behaviour is enhanced when the EQ mols are between 1.5 and 3 (Ex A, A', B, E, E', F, F').

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Formulations containing the composition of the present invention are exemplified by the following:

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The detergent compositions of the present invention may be formulated as shampoos, baby shampoos, conditioning shampoos, bath gels, hair conditioners, for manual dishwashing, and as all purpose cleaners which are exemplified below (all values indicated are weight percentages):

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Baby Shampoo

COMPONENTS	BS1	BS2	
Deionized water	to 100	to 100	55
Sodium Lauryl sulfate (27% Dry) (Emal @ 227E from Kao)	25.0	8.0	
Sodium Cocoamphoacetate (40% Dry) (Betadet @ SHC-2 from Kao)	7.5	15.0	
Example A product	2.0	2.0	
Lauryl hydroxysultaine (45% Dry) (Betadet @ S-20 from Kao)	4.0		60
PEG-20 Sorbitan Laurate (Kaopan @ TW-L-120 from Kao)	—	1.0	
PEG-120 Methylglucose dioleate (Glucamate-DOE-120 @ from Amercabol)	—	0.2	
NaCl	0.2	—	65
Preservative	0.05	0.05	

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-continued

		<u>Baby Shampoo</u>	
30	COMPONENTS	BS1	BS2
<u>ANALYSIS</u>			
35	Appearance	Transparent viscous liquid	Transparent viscous liquid
pH (100%)	6.5-7.5	6.5-7.5	
Viscosity (cps) 20° C.	5000-7000	1000-2000	
Turbidity point (° C.)	<0	<0	
% Dry matter	12.5-14.5	10.5-12.5	
Stability	OK	OK	

45	COMPONENTS	
<u>Shampoo</u>		

50	Deionized water	to 100
	Sodium Lauryl sulfate (70% Dry) (Emal ® 270E from Kao)	23.0
	Cocoamidopropoxybetaine (48% Dry) (Betadet ® HR-50K from Kao)	10.0
	Example B product	1.8
	Pearling agent (Danox ® P-15 from Kao)	3.0
	Perfume	e.q.
55	NaCl	e.q.
	Preservative	e.q.
<u>ANALYSIS</u>		
	Appearance	Pearled viscous liquid
60	pH (100%)	6.0-7.0
	Viscosity (cps) 20° C.	=8000
	% Dry matter	24-26
	Stability	OK

65	Conditioning shampoo	
	Deionized water	to 100
	Sodium Lauryl sulfate (27% Dry)	32.0

-continued

COMPONENTS

(Emal ® 227E from Kao)	
Sodium Cocoamphoacetate (40% Dry)	7.5
(Betadet ® SHC-2 from Kao)	
Example E product	3.5
Lauryl hydroxysultaine (45% Dry)	5.0
(Betadet ® S-20 from Kao)	
Oleic esterquat (80% Dry Matter)	0.5
(Tetraethyl ® CO-40 from Kao)	
Pearling agent (Danox ® BF-22 from Kao)	3.0
Perfume	e.q.
NaCl	e.q.
Preservative	e.q.

ANALYSIS

Appearance	Pearled viscous liquid
pH (100%)	6.0-6.5
Viscosity (cps) 20° C.	≈7000
% Dry matter	19-21
Stability	OK

Bath gelCOMPONENTS

Deionized water	to 100
Sodium Lauryl sulfate (27% Dry)	37.0
(Emal ® 277 E from Kao)	
Cocoamidopropoxybetaine (34% Dry)	10.0
(Betadet ® HR from Kao)	
Example F product	2.5
Perfume	0.5
NaCl	0.5
Preservative: Kathon CG ® from Rohm & Haas	0.05
EDTA.Na ₂	0.05

ANALYSIS

Appearance	Transparent viscous liquid
pH (100%)	5.0-6.0
Viscosity (cps) 20° C.	6000-8000
Turbidity point (° C.)	<0
% Dry matter	18-20
Stability	OK

Hair conditionerCOMPONENTS

	HC1	HC2
Deionized water	to 100	to 100
Propylene glycol	2.0	2.0
Dioleic esterquat (80% Dry Matter) (Tetraethyl ® CO-40 from Kao)	1.9	—
Cetrimonium Chloride (25% Dry) (Quartamin ® 60W25 from Kao)	—	6.0
Cetearyl alcohol (Kalcot ® 6870 from Kao)	3.0	3.0
Example A product	0.5	0.5
Perfume	e.q.	e.q.
Preservative	e.q.	e.q.

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